AGTCCCAGACGGCTTTTCCCAGAGAGCTAAAAGAGAAGGCCCAGAGAATGTCCTCCCAG 5 CCAGCAGGAACCAGACCTCCCCGGGGCCACAGAGGACTACTCCTATGGCAGCTGGTAC ACCAGCATACCACCCGGCCTGTACCACGCCTGCCTGGCCTCGCTGTCAATCCTTGTGCTG CTGCTCCTGGCCATGCTGAGGCGCCGCCAGCTCTGGCCTGACTGTGCGTGGCAGG CCCGGCCTGCCAGCCCTGTGGATTTCTTGGCTGGGGACAGGCCCCGGGCAGTGCCTGCT 10 GCTGTTTTCATGGTCCTCCTGAGCTCCCTGTGTTTGCTGCTCCCCGACGAGGACGCATTG CCCTTCCTGACTCTCGCCTCAGCACCCAGCCAAGATGGGAAAACTGAGGCTCCAAGAGGG GCCTGGAAGATACTGGGACTGTTCTATTATGCTGCCCTCTACTACCCTCTGGCTGCCTGT GCCACGGCTGGCCACACAGCTGCACACCTGCTCGGCAGCACGCTGTCCTGGGCCCACCTT GGGGTCCAGGTCTGGCAGAGGGCAGAGTGTCCCCAGGTGCCCAAGATCTACAAGTACTAC 15 TCCCTGCTGGCCTCCCTGCTCCTGCTGGGCCTCGGATTCCTGAGCCTTTGGTACCCT GTGCAGCTGGTGAGAAGCTTCAGCCGTAGGACAGGAGCAGGCTCCAAGGGGCTGCAGAGC AGCTACTCTGAGGAATATCTGAGGAACCTCCTTTGCAGGAAGAAGCTGGGAAGCAGCTAC CACACCTCCAAGCATGGCTTCCTGTCCTGGGCCCGCGTCTGCTTGAGACACTGCATCTAC ACTCCACAGCCAGGATTCCATCTCCCGCTGAAGCTGGTGCTTTCAGCTACACTGACAGGG ₫20 ACGGCCATTTACCAGGTGGCCCTGCTGCTGCTGGTGGGCGTGGTACCCACTATCCAGAAG GTGAGGGCAGGGGTCACCACGGATGTCTCCTACCTGCTGGCCGGCTTTGGAATCGTGCTC TCCGAGGACAAGCAGGAGGTGGTGGAGCTGGTGAAGCACCATCTGTGGGCTCTGGAAGTG TGCTACATCTCAGCCTTGGTCTTGTCCTGCTTACTCACCTTCCTGGTCCTGATGCGCTCA CTGGTGACACAGGACCAACCTTCGAGCTCTGCACCGAGGAGCTGCCCTGGACTTGAGT <u>1</u>25 CCCTTGCATCGGAGTCCCCATCCCTCCCGCCAAGCCATATTCTGTTGGATGAGCTTCAGT GCCTACCAGACAGCCTTTATCTGCCTTGGGCTCCTGGTGCAGCAGATCATCTTCCTG GGAACCACGGCCTGGCCTTCCTGGTGCTCATGCCTGTGCTCCATGGCAGGAACCTCCTG CTCTTCCGTTCCCTGGAGTCCTCGTGGCCCTTCTGGCTGACTTTGGCCCTGGCTGTGATC CTGCAGAACATGGCAGCCCATTGGGTCTTCCTGGAGACTCATGATGGACACCCACAGCTG **54** 3 0 ACCAACCGGCGAGTGCTCTATGCAGCCACCTTTCTTCTCTCCCCCTCAATGTGCTGGTG GGTGCCATGGTGGCCACCTGGCGAGTGCTCCTCTCTGCCCTCTACAACGCCATCCACCTT GGCCAGATGGACCTCAGCCTGCTGCCACCGAGAGCCGCCACTCTCGACCCCGGCTACTAC ACGTACCGAAACTTCTTGAAGATTGAAGTCAGCCAGTCGCATCCAGCCATGACAGCCTTC TGCTCCTGCTCCTGCAAGCGCAGAGCCTCCTACCCAGGACCATGGCAGCCCCCCAGGAC 35 AGCCTCAGACCAGGGGAGGAAGACGAAGGGATGCAGCTGCTACAGACAAAGGACTCCATG GCCAAGGGAGCTAGGCCCGGGGCCAGCCGCGGCAGGGCTCGCTGGGGTCTGGCCTACACG CTGCTGCACAACCCAACCCTGCAGGTCTTCCGCAAGACGGCCCTGTTGGGTGCCAATGGT GCCCAGCCCTGAGGGCAGGGAAGGTCAACCCACCTGCCCATCTGTGCTGAGGCATGTTCC TGCCTACCATCCTCCCCCCGGCTCTCCCCAGCATCACACCAGCCATGCAGCCA 40 GGCTCTGCTCCACCCACTTGGCTATGGGAGAGCCAGCAGGGGTTCTGGAGAAAAAACTG GTGGGTTAGGGCCTTGGTCCAGGAGCCAGTTGAGCCAGGGCAGCCACATCCAGGCGTCTC CCTACCCTGGCTCTGCCATCAGCCTTGAAGGGCCTCGATGAAGCCTTCTCTGGAACCACT CCAGCCCAGCTCCACCTTGGCCTTCACGCTGTGGAAGCAGCCAAGGCACTTCCT 45 CACCCCTCAGCGCCCACGGACCTCTCTGGGGAGTGGCCGGAAAGCTCCCGGTCCTCTGGC CTGCAGGGCAGCCCAAGTCATGACTCAGACCAGGTCCCACACTGAGCTGCCCACACTCGA GAGCCAGATATTTTTGTAGTTTTTATGCCTTTTGGCTATTATGAAAGAGGTTAGTGTTTC 

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MSSQPAGNQTSPGATEDYSYGSWYIDEPQGGEELQPEGEVPSCHTSIPPGLYHACLASLS ILVLLLLAMLVRRRQLWPDCVRGRPGLPSPVDFLAGDRPRAVPAAVFMVLLSSLCLLLPD EDALPFLTLASAPSQDGKTEAPRGAWKILGLFYYAALYYPLAACATAGHTAAHLLGSTLS WAHLGVQVWQRAECPQVPKIYKYYSLLASLPLLLGLGFLSLWYPVQLVRSFSRRTGAGSK GLQSSYSEEYLRNLLCRKKLGSSYHTSKHGFLSWARVCLRHCIYTPQPGFHLPLKLVLSA TLTGTAIYQVALLLLVGVVPTIQKVRAGVTTDVSYLLAGFGIVLSEDKQEVVELVKHHLW ALEVCYISALVLSCLLTFLVLMRSLVTHRTNLRALHRGAALDLSPLHRSPHPSRQAIFCW MSFSAYQTAFICLGLLVQQIIFFLGTTALAFLVLMPVLHGRNLLLFRSLESSWPFWLTLA LAVILQNMAAHWVFLETHDGHPQLTNRRVLYAATFLLFPLNVLVGAMVATWRVLLSALYN AIHLGQMDLSLLPPRAATLDPGYYTYRNFLKIEVSQSHPAMTAFCSLLLQAQSLLPRTMA APQDSLRPGEEDEGMQLLQTKDSMAKGARPGASRGRARWGLAYTLLHNPTLQVFRKTALL GANGAQP

Important features of the protein:

15 Signal peptide:

5

10

55

132-140

```
None
Transmembrane domain:
ū
   20
[]
        54-69
        102-119
į.a
        148-166
ſ.
         207-222
ļ.
   25
         301-320
TU
         364-380
Ŧ
         431-451
474-489
الم ا
         560-535
ļ4
   30
[]
         Motif file:
         Motif name: N-glycosylation site.
ļA
               8-12
   35
         Motif name: N-myristoylation site.
              50-56
             176-182
             241-247
   40
             317-323
             341-347
             525-531
             627-633
   45
             631-637
             640-646
             661-667
         Motif name: Prokaryotic membrane lipoprotein lipid attachment site.
   50
             364-375
         Motif name: ATP/GTP-binding site motif A (P-loop).
```

### 

### FIGURE 3A

PRO

XXXXXXXXXXXXXXX

(Length = 15 amino acids)

Comparison Protein

XXXXXYYYYYYY

(Length = 12 amino acids)

5

10

% amino acid sequence identity =

(the number of identically matching amino acid residues between the two polypeptide sequences as determined by ALIGN-2) divided by (the total number of amino acid residues of the PRO polypeptide) =

5 divided by 15 = 33.3%

### 

### FIGURE 3B

**PRO** 

XXXXXXXXX

(Length = 10 amino acids)

Comparison Protein

XXXXXYYYYYYZZYZ

(Length = 15 amino acids)

5

10

% amino acid sequence identity =

(the number of identically matching amino acid residues between the two polypeptide sequences as determined by ALIGN-2) divided by (the total number of amino acid residues of the PRO polypeptide) =

5 divided by 10 = 50%

### FIGURE 3C

PRO-DNA	ининининини	(Length	=	14
nucleotides)				
Comparison DNA	NNNNNLLLLLLLLLL	(Length	=	16
nucleotides)				

% nucleic acid sequence identity =

(the number of identically matching nucleotides between the two nucleic acid sequences as determined by ALIGN-2) divided by (the total number of nucleotides of the PRO-DNA nucleic acid sequence) =

6 divided by 14 = 42.9%

15

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5

### FIGURE 3D

PRO-DNA

ииииииииии

(Length = 12 nucleotides)

Comparison DNA

NNNNLLLVV

(Length = 9 nucleotides)

5

10

% nucleic acid sequence identity =

(the number of identically matching nucleotides between the two nucleic acid sequences as determined by ALIGN-2) divided by (the total number of nucleotides of the PRO-DNA nucleic acid sequence) =

4 divided by 12 = 33.3%

### FIGURE 4A

```
* C-C increased from 12 to 15
     5
            * Z is average of EQ
            * B is average of ND
            * match with stop is M; stop-stop = 0; J (joker) match = 0
           #define M
                             -8
                                      /* value of a match with a stop */
   10
                     day[26][26] = {
                  ABCDEFGHIJKLMNOPQRSTUVWXYZ*/
           /*
           /* A */
                      {2, 0, -2, 0, 0, -4, 1, -1, -1, 0, -1, -2, -1, 0, M, 1, 0, -2, 1, 1, 0, 0, -6, 0, -3, 0},
           /* B */
                      { 0, 3,-4, 3, 2,-5, 0, 1,-2, 0, 0,-3,-2, 2, M,-1, 1, 0, 0, 0, 0,-2,-5, 0,-3, 1},
                      \{-2,-4,15,-5,-5,-4,-3,-3,-2,0,-5,-6,-5,-4,\_M,-3,-5,-4,0,-2,0,-2,-8,0,0,-5\},
   15
           /* C */
           /* D */
                      { 0, 3,-5, 4, 3,-6, 1, 1,-2, 0, 0,-4,-3, 2, M,-1, 2,-1, 0, 0, 0,-2,-7, 0,-4, 2},
           /* E */
                      \{0, 2, -5, 3, 4, -5, 0, 1, -2, 0, 0, -3, -2, 1, M, -1, 2, -1, 0, 0, 0, -2, -7, 0, -4, 3\},\
           /* F */
                      \{-4, -5, -4, -6, -5, 9, -5, -2, 1, 0, -5, 2, 0, -4, M, -5, -5, -4, -3, -3, 0, -1, 0, 0, 7, -5\},\
           /* G */
                      {1, 0,-3, 1, 0,-5, 5,-2,-3, 0,-2,-4,-3, 0, M,-1,-1,-3, 1, 0, 0,-1,-7, 0,-5, 0},
   20
           /* H */
                      {-1, 1,-3, 1, 1,-2,-2, 6,-2, 0, 0,-2,-2, 2, M, 0, 3, 2,-1,-1, 0,-2,-3, 0, 0, 2},
           /* 1 */
                      \{-1, -2, -2, -2, -2, 1, -3, -2, 5, 0, -2, 2, 2, 2, -2, M, -2, -2, -1, 0, 0, 4, -5, 0, -1, -2\},\
                      /* J */
/* K */
                      {-1, 0,-5, 0, 0,-5,-2, 0,-2, 0, 5,-3, 0, 1, M,-1, 1, 3, 0, 0, 0,-2,-3, 0,-4, 0},
           /* L */
                      {-2,-3,-6,-4,-3, 2,-4,-2, 2, 0,-3, 6, 4,-3,_M,-3,-2,-3,-1, 0, 2,-2, 0,-1,-2},
   25
           /* M */
                      \{-1,-2,-5,-3,-2,0,-3,-2,2,0,0,4,6,-2,M,-2,-1,0,-2,-1,0,2,-4,0,-2,-1\}
           /* N */
                      { 0, 2,-4, 2, 1,-4, 0, 2,-2, 0, 1,-3,-2, 2, M,-1, 1, 0, 1, 0, 0,-2,-4, 0,-2, 1}.
Ļá
           /* O */
                                                               d
           0, M, M},
           /* P */
                      { 1,-1,-3,-1,-1,-5,-1, 0,-2, 0,-1,-3,-2,-1, M, 6, 0, 0, 1, 0, 0,-1,-6, 0,-5, 0},
řá
   30
           /* Q */
                      { 0, 1,-5, 2, 2,-5,-1, 3,-2, 0, 1,-2,-1, 1, M, 0, 4, 1,-1,-1, 0,-2,-5, 0,-4, 3},
fU
           /* R */
                      {-2, 0,-4,-1,-1,-4,-3, 2,-2, 0, 3,-3, 0, 0, M, 0, 1, 6, 0,-1, 0,-2, 2, 0,-4, 0},
æ
                      { 1, 0, 0, 0, 0, -3, 1, -1, -1, 0, 0, -3, -2, 1, M, 1, -1, 0, 2, 1, 0, -1, -2, 0, -3, 0},
           /* S */
[.]
           /* T */
                      { 1, 0,-2, 0, 0,-3, 0,-1, 0, 0, 0,-1,-1, 0, M, 0,-1,-1, 1, 3, 0, 0,-5, 0,-3, 0},
                      /* U */
   35
           /* V */
                      { 0,-2,-2,-2,-1,-1,-2, 4, 0,-2, 2, 2,-2, M,-1,-2,-2,-1, 0, 0, 4,-6, 0,-2,-2},
           /* W */
                      {-6,-5,-8,-7,-7, 0,-7,-3,-5, 0,-3,-2,-4,-4, M,-6,-5, 2,-2,-5, 0,-6,17, 0, 0,-6},
[]
           /* X */
                      /* Y */
                      {-3,-3, 0,-4,-4, 7,-5, 0,-1, 0,-4,-1,-2,-2, M,-5,-4,-4,-3,-3, 0,-2, 0, 0,10,-4},
           1* Z *1
                      { 0, 1,-5, 2, 3,-5, 0, 2,-2, 0, 0,-2,-1, 1, M, 0, 3, 0, 0, 0, 0, 0,-2,-6, 0,-4, 4}
   40
           };
```

45

50

### FIGURE 4B

```
*/
            #include < stdio.h>
      5
            #include < ctype.h>
            #define MAXJMP
                                         16
                                                  /* max jumps in a diag */
            #define MAXGAP
                                         24
                                                  /* don't continue to penalize gaps larger than this */
            #define JMPS
                                         1024
                                                  /* max jmps in an path */
    10
            #define MX
                                         4
                                                  /* save if there's at least MX-1 bases since last jmp */
            #define DMAT
                                         3
                                                  /* value of matching bases */
            #define DMIS
                                         0
                                                  /* penalty for mismatched bases */
                                         8
            #define DINSO
                                                  /* penalty for a gap */
    15
            #define DINS1
                                         Ì
                                                  /* penalty per base */
            #define PINSO
                                         8
                                                  /* penalty for a gap */
            #define PINS1
                                                  /* penalty per residue */
            struct jmp {
    20
                                         n[MAXJMP];
                                                            /* size of jmp (neg for dely) */
                      short
                                         x[MAXJMP];
                                                            /* base no. of jmp in seq x */
                      unsigned short
/* limits seq to 2°16 -1 */
            };
            struct diag {
    25
                                                            /* score at last jmp */
                      int
                                         score;
                      long
                                         offset;
                                                            /* offset of prev block */
ļā
                                                            /* current jmp index */
                      short
                                         ijmp;
O
                                                            /* list of jmps */
                      struct jmp
                                         jp;
ļ.š
            };
   30
TU
            struct path {
Ð
                                                  /* number of leading spaces */
                     int
n[JMPS];/* size of jmp (gap) */
                      short
                               x[JMPS];/* loc of jmp (last elem before gap) */
                     int
<u>14</u> 35
            };
char
                               *ofile;
                                                            /* output file name */
D
                               *namex[2];
                                                            /* seq names: getseqs() */
            char
Ĺá
                               *prog;
                                                            /* prog name for err msgs */
            char
    40
                               *seqx[2];
                                                            /* seqs: getseqs() */
            char
                               dmax;
                                                            /* best diag: nw() */
            int
            int
                               dmax0;
                                                            /* final diag */
                                                            /* set if dna: main() */
            int
                               dna;
                                                            /* set if penalizing end gaps */
            int
                               endgaps;
   45
                                                            /* total gaps in seqs */
            int
                               gapx, gapy;
                                                            /* seq lens */
            int
                               len0, len1;
                                                            /* total size of gaps */
            int
                               ngapx, ngapy;
            int
                               smax;
                                                            /* max score: nw() */
            int
                               *xbm;
                                                            /* bitmap for matching */
   50
                               offset;
                                                            /* current offset in jmp file */
            long
                               *dx;
                                                            /* holds diagonals */
            struct
                     diag
            struct
                     path
                                                            /* holds path for seqs */
                               pp[2];
            char
                               *calloc(), *malloc(), *index(), *strcpy();
   55
            char
                               *getseq(), *g_calloc();
```

### FIGURE 4C

```
/* Needleman-Wunsch alignment program
              usage: progs file1 file2
     5
                where file1 and file2 are two dna or two protein sequences.
                The sequences can be in upper- or lower-case an may contain ambiguity
               Any lines beginning with ';', '>' or '<' are ignored
               Max file length is 65535 (limited by unsigned short x in the jmp struct)
               A sequence with 1/3 or more of its elements ACGTU is assumed to be DNA
    10
                Output is in the file "align.out"
             * The program may create a tmp file in /tmp to hold info about traceback.
             * Original version developed under BSD 4.3 on a vax 8650
    15
            #include "nw.h"
            #include "day.h"
            static
                       dbval[26] = {
                      1,14,2,13,0,0,4,11,0,0,12,0,3,15,0,0,0,5,6,8,8,7,9,0,10,0
20
            };
٠...
الله
                      _{pbval[26]} = {
            static
                      \overline{1}, 2[(1 < < ('D'-'A'))](1 < < ('N'-'A')), 4, 8, 16, 32, 64,
[]
                      128, 256, 0xFFFFFFF, 1 < < 10, 1 < < 11, 1 < < 12, 1 < < 13, 1 < < 14,
<u>ļ</u>4
                      1 < < 15, 1 < < 16, 1 < < 17, 1 < < 18, 1 < < 19, 1 < < 20, 1 < < 21, 1 < < 22.
    25
ſΰ
                      1 < < 23, 1 < < 24, 1 < < 25  \{(1 < < ('E'-'A'))\}(1 < < ('Q'-'A'))
ļá
            };
fIJ
                                                                                                                                 main
            main(ac, av)
35
   30
                      int
                                ac;
                                *av[];
                      char
الم
į d
                      prog = av[0];
[]
[] 35
                      if (ac != 3) {
                                fprintf(stderr, "usage: %s file1 file2\n", prog);
                                fprintf(stderr, "where file1 and file2 are two dna or two protein sequences.\n");
ļā
                                fprintf(stderr, "The sequences can be in upper- or lower-case\n");
                                fprintf(stderr, "Any lines beginning with '; ' or ' < ' are ignored\n");
                                fprintf(stderr, "Output is in the file \"align.out\"\n");
    40
                                exit(1);
                      }
                      namex[0] = av[1];
                      namex[1] = av[2];
                      seqx[0] = getseq(namex[0], \&len0);
    45
                      seqx[1] = getseq(namex[1], &len1);
                      xbm = (dna)? _dbval : _pbval;
                                                              /* 1 to penalize endgaps */
                      endgaps = 0;
                                                             /* output file */
                      ofile = "align.out";
    50
                                          /* fill in the matrix, get the possible jmps */
                      nw();
                                          /* get the actual jmps */
                      readjmps();
                                          /* print stats, alignment */
                      print();
    55
                      cleanup(0);
                                          /* unlink any tmp files */
```

### FIGURE 4D

```
/* do the alignment, return best score: main()
            * dna: values in Fitch and Smith, PNAS, 80, 1382-1386, 1983
            * pro: PAM 250 values
            * When scores are equal, we prefer mismatches to any gap, prefer
            * a new gap to extending an ongoing gap, and prefer a gap in seqx
            * to a gap in seq y.
            nw()
                                                                                                                                  nw
   10
            {
                                                            /* seqs and ptrs */
                     char
                                         *px, *py;
                                                            /* keep track of dely */
                     int
                                         *ndely, *dely;
                     int
                                         ndelx, delx;
                                                            /* keep track of delx */
                                         *tmp;
                                                            /* for swapping row0, row1 */
                     int
   15
                                                            /* score for each type */
                     int
                                         mis;
                     int
                                         ins0, ins1;
                                                            /* insertion penalties */
                                                            /* diagonal index */
                     register
                                         id;
                     register
                                                            /* jmp index */
                                         ij;
                                         *col0, *col1;
                                                            /* score for curr, last row */
                     register
   20
                     register
                                                            /* index into seqs */
                                         xx, yy;
1
1
1
1
2
5
                      dx = (struct diag *)g_calloc("to get diags", len0+len1+1, sizeof(struct diag));
                     ndely = (int *)g calloc("to get ndely", len1+1, sizeof(int));
                      dely = (int *)g calloc("to get dely", len1 + 1, sizeof(int));
Ļ
                      col0 = (int *)g calloc("to get col0", len1+1, sizeof(int));
ťÔ
                      coll = (int *)g calloc("to get coll", len1 + 1, sizeof(int));
                      ins0 = (dna)? DINS0 : PINS0;
ķ
                      ins1 = (dna)? DINS1 : PINS1;
N 30
                      smax = -10000;
17
                     if (endgaps) {
                               for (col0[0] = dely[0] = -ins0, yy = 1; yy < = len1; yy + +) {
۱.,[
                                         col0[yy] = dely[yy] = col0[yy-1] - ins1;
14
[]
   35
                                         ndely[yy] = yy;
/* Waterman Bull Math Biol 84 */
                               col0[0] = 0;
<u>}</u> .4
                     }
                     else
   40
                               for (yy = 1; yy < = len1; yy++)
                                         dely[yy] = -ins0;
                     /* fill in match matrix
   45
                     for (px = seqx[0], xx = 1; xx <= len0; px++, xx++) {
                               /* initialize first entry in col
                               */
                               if (endgaps) {
                                        if (xx = = 1)
   50
                                                   col1[0] = delx = -(ins0 + ins1);
                                                  col1[0] = delx = col0[0] - ins1;
                                         ndelx = xx;
   55
                               else {
                                        col1[0] = 0;
                                        delx = -ins0;
                                        ndelx = 0;
                               }
```

### FIGURE 4E

for  $(py = seqx[1], yy = 1; yy <= len1; py++, yy++) {$ mis = col0[yy-1];5 if (dna) mis += (xbm[\*px-'A']&xbm[\*py-'A'])? DMAT : DMIS;else  $mis + = _day[*px-'A'][*py-'A'];$ /\* update penalty for del in x seq; 10 \* favor new del over ongong del \* ignore MAXGAP if weighting endgaps if (endgaps | | ndely[yy] < MAXGAP) { if (col0[yy] - ins0 > = dely[yy]) { 15 dely[yy] = col0[yy] - (ins0 + ins1);ndely[yy] = 1;} else { dely[yy] = insl;ndely[yy] + +;20 } else { if  $(col0[yy] - (ins0 + ins1) > = dely[yy]) {$ dely[yy] = col0[yy] - (ins0 + ins1);ndely[yy] = 1;25 } else ĹŪ ndely[yy] + +; ļ. } /\* update penalty for del in y seg; 30 \* favor new del over ongong del if (endgaps || ndelx < MAXGAP) { if  $(coll[yy-1] - ins0 > = delx) {$ ļå **1** 35 delx = coll[yy-1] - (ins0 + ins1);ndelx = 1;} else { ļá delx -= insl;ndelx + +;} 40 } else { if  $(coll[yy-1] - (ins0 + ins1) > = delx) {$ delx = coll[yy-1] - (ins0 + ins1);ndelx = 1;45 } else ndelx++; } /\* pick the maximum score; we're favoring \* mis over any del and delx over dely 50

60

55

...nw

### FIGURE 4F

...pw

```
id = xx - yy + lenl - 1;
                                      if (mis > = delx && mis > = dely(yy))
                                               coll[yy] = mis;
    5
                                      else if (delx > = dely(yy)) {
                                               coll[yy] = delx;
                                               ij = dx[id].ijmp;
                                               if (dx[id].jp.n[0] && (!dna | | | (ndelx > = MAXJMP)
                                               && xx > dx[id].jp.x[ij]+MX) || mis > dx[id].score+DINS0)) {
  10
                                                         dx[id].ijmp++;
                                                         if (++ij > = MAXJMP) {
                                                                  writejmps(id);
                                                                  ij = dx{id}.ijmp = 0;
                                                                  dx[id].offset = offset;
   15
                                                                  offset += sizeof(struct jmp) + sizeof(offset);
                                               dx[id].jp.n[ij] = ndelx;
                                                dx[id].jp.x[ij] = xx;
   20 .
                                                dx[id].score = delx;
[]
                                      else {
Ū
₩ 25
                                                coll[yy] = dely[yy];
                                                ij = dx[id].ijmp;
                    if (dx[id].jp.n[0] && (!dna || (ndely[yy] > = MAXJMP
ļä
                                                && xx > dx[id].jp.x[ij]+MX) || mis > dx[id].score+D1NS0)) {
[A
                                                         dx[id].ijmp++;
                                                         if (++ij > = MAXJMP) {
   30
                                                                   writejmps(id);
                                                                   ij = dx[id].ijmp = 0;
=
                                                                   dx[id].offset = offset;
1
                                                                   offset += sizeof(struct jmp) + sizeof(offset);
    35
dx[id].jp.n[ij] = -ndely[yy];
                                                dx[id].jp.x[ij] = xx;
                                                dx[id].score = dely[yy];
    40
                                       if (xx =   len0 && yy <  len1) {
                                                 /* last col
                                                 */
                                                 if (endgaps)
                                                          coll[yy] = ins0 + ins1*(len1-yy);
    45
                                                 if (coll[yy] > smax) {
                                                          smax = coll[yy];
                                                          dmax = id;
                                                 }
    50
                              if (endgaps && xx < len0)
                                        coll[yy-1] -= ins0 + ins1*(len0-xx);
                               if (coll[yy-1] > smax) {
                                        smax = coll[yy-1];
     55
                                        dmax = id;
                               tmp = col0; col0 = col1; col1 = tmp;
                      (void) free((char *)ndely);
     60
                      (void) free((char *)dely);
                                                                                       Page 4 of nw.c
                      (void) free((char *)col0);(void) free((char *)col1);}
```

### FIGURE 4G

```
print() -- only routine visible outside this module
     5
            * getmat() -- trace back best path, count matches: print()
            * pr align() -- print alignment of described in array p[]: print()
            * dumpblock() -- dump a block of lines with numbers, stars: pr_align()
   10
            * nums() - put out a number line: dumpblock()
            * putline() -- put out a line (name, [num], seq, [num]): dumpblock()
            * stars() - -put a line of stars: dumpblock()
            * stripname() -- strip any path and prefix from a seqname
            */
   15
            #include "nw.h"
            #define SPC
            #define P LINE
                              256
                                        /* maximum output line */
            #define P SPC
                                        /* space between name or num and seq */
   20
                               3
                      day[26][26];
            extern
/* set output line length */
            int
                     olen;
            FILE
                      *fx:
                                         /* output file */
   25
                                                                                                                               print
            print()
                     int
                               Ix, ly, firstgap, lastgap;
                                                            /* overlap */
Ü
1 4 3 0
                     if ((fx = fopen(ofile, "w")) = = 0) {
TU
                               fprintf(stderr," %s: can't write %s\n", prog, ofile);
                               cleanup(1);
=
fprintf(fx, " < first sequence: %s (length = %d)\n", namex[0], len0);
                     fprintf(fx, " < second sequence: %s (length = %d)\n", namex[1], len1);
40
                     olen = 60;
                     lx = len0;
                     ly = len1;
                      firstgap = lastgap = 0;
                      if (dmax < len1 - 1) {
                                                  /* leading gap in x */
                               pp[0].spc = firstgap = len1 - dmax - 1;
                               ly -= pp{0}.spc;
                     else if (dmax > len1 - 1) { /* leading gap in y */
   45
                               pp[1].spc = firstgap = dmax - (len1 - 1);
                               lx -= pp[1].spc;
                      if (dmax0 < len0 - 1) {
                                                  /* trailing gap in x */
                               lastgap = len0 - dmax0 - 1;
   50
                               lx -= lastgap;
                      else if (dmax0' > len0 - 1) { /* trailing gap in y */
                               lastgap = dmax0 - (len0 - 1);
                               ly -= lastgap;
    55
                      getmat(lx, ly, firstgap, lastgap);
                      pr_align();
            }
```

### FIGURE 4H

```
* trace back the best path, count matches
            */
     5
           static
                                                                                                                      getmat
           getmat(lx, ly, firstgap, lastgap)
                                                         /* "core" (minus endgaps) */
                    int
                             lx, ly;
                                                         /* leading trailing overlap */
                             firstgap, lastgap;
                    int
           {
                                       nm, i0, i1, siz0, siz1;
   10
                    int
                                       outx[32];
                     char
                                       pct;
                     double
                                       n0, n1;
                     register
                                       *p0, *p1;
                     register char
    15
                     /* get total matches, score
                     i0 = i1 = siz0 = siz1 = 0;
                     p0 = seqx[0] + pp[1].spc;
                     p1 = seqx[1] + pp[0].spc;
    20
                     n0 = pp[1].spc + 1;
                     n1 = pp[0].spc + 1;
Ę
                     nm = 0;
1
                     while (*p0 && *p1) {
   25
                              if (siz0) {
                                       pl++;
                                       nl++;
to
                                       siz0--;
ļå
                              }
    30
                              else if (siz1) {
ſIJ
                                       p0++;
=
                                       n0++;
IJ
                                       sizl--;
    35
                              }
                              else {
if (xbm[*p0-'A']&xbm[*p1-'A'])
                                                 nm++;
                                        if (n0++=pp\{0\}.x[i0])
ļ#
                                                 siz0 = pp[0].n[i0++];
    40
                                        if(n1++==pp[1].x[i1])
                                                 siz1 = pp[1].n[i1++];
                                        p0++;
                                        pl++;
    45
                               }
                      }
                      /* pct homology:
                      * if penalizing endgaps, base is the shorter seq
                      * else, knock off overhangs and take shorter core
     50
                      */
                      if (endgaps)
                               1x = (len0 < len1)? len0 : len1;
                      else
     55
                               lx = (lx < ly)? lx : ly;
                      pct = 100.*(double)nm/(double)lx;
                      fprintf(fx, "\n");
                      fprintf(fx, " < %d match%s in an overlap of %d: %.2f percent similarity\n",
                               nm, (nm = = 1)? "" : "es", lx, pct);
     60
```

```
...getmat
                     fprintf(fx, " < gaps in first sequence: %d", gapx);
                    if (gapx) {
                              (void) sprintf(outx, " (%d %s%s)",
    5
                                       ngapx, (dna)? "base": "residue", (ngapx == 1)? "": "s");
                              fprintf(fx,"%s", outx);
                     fprintf(fx, ", gaps in second sequence: %d", gapy);
   10
                     if (gapy) {
                              (void) sprintf(outx, " (%d %s%s)",
                                       ngapy, (dna)? "base": "residue", (ngapy == 1)? "": "s");
                              fprintf(fx, "%s", outx);
                     }
   15
                     if (dna)
                               fprintf(fx,
                               "n < score: %d (match = %d, mismatch = %d, gap penalty = %d + %d per base)n",
                              smax, DMAT, DMIS, DINSO, DINSI);
                     else
   20
                               fprintf(fx,
                               "\n < score: %d (Dayhoff PAM 250 matrix, gap penalty = %d + %d per residue)\n",
                               smax, PINSO, PINS1);
if (endgaps)
                               fprintf(fx,
                               " < endgaps penalized. left endgap: %d %s%s, right endgap: %d %s%s\n",
10
   25
                               firstgap, (dna)? "base": "residue", (firstgap = = 1)? "": "s",
                               lastgap, (dna)? "base": "residue", (lastgap = = 1)? "": "s");
ŀå
                     else
đ
                               fprintf(fx, " < endgaps not penalized\n");
<u>1</u> 30
            }
TU
                                                  /* matches in core -- for checking */
             static
                               nm;
Ξ
                                                  /* lengths of stripped file names */
             static
                               lmax;
[]
                                                  /* imp index for a path */
                               ij[2];
             static
                                                  /* number at start of current line */
   35
             static
                               nc[2];
                                                  /* current elem number -- for gapping */
ŀ≖
                               ni[2];
             static
13
             static
                               siz[2];
                                                  /* ptr to current element */
static char
                               *ps[2];
                                                  /* ptr to next output char slot */
             static char
                               *po[2];
                               out[2][P_LINE]; /* output line */
    40
             static char
                               star[P LINE];
                                                  /* set by stars() */
             static char
             * print alignment of described in struct path pp[]
             */
    45
            static
                                                                                                                         pr align
            pr_align()
            {
                                                  /* char count */
                                         nn;
                      int
    50
                      int
                                         more;
                      register
                                         i;
                      for (i = 0, lmax = 0; i < 2; i++)
                               nn = stripname(namex[i]);
    55
                               if (nn > lmax)
                                         lmax = nn;
                               nc[i] = 1;
                                ni[i] = 1;
                                siz[i] = ij[i] = 0;
    60
                                ps[i] = seqx[i];
                                                                                                  Page 3 of nwprint.c
                                po[i] = out[i];
```

### FIGURE 4J

```
...pr align
                     for (nn = nm = 0, more = 1; more;)
                              for (i = more = 0; i < 2; i++) {
     5
                                        * do we have more of this sequence?
                                        */
                                       if (!*ps[i])
                                                continue;
   10
                                       more++;
                                                         /* leading space */
                                       if (pp[i].spc) {
                                                *po[i]++='';
   15
                                                pp[i].spc--;
                                       }
                                       else if (siz[i]) {
                                                       /* in a gap */
                                                 *po[i] + + = '-';
                                                siz[i]--;
   20
                                       }
                                       else {
                                                          /* we're putting a seq element
*po[i] = *ps[i];
                                                if (islower('ps[i]))
                                                          *ps[i] = toupper(*ps[i]);
                                                ps[i]++;
ίū
14 30
                                                 * are we at next gap for this seq?
TL.
                                                if (ni[i] = pp[i].x[ij[i]]) {
                                                           * we need to merge all gaps
<sup>≒</sup>₄ 35
                                                           * at this location
ļå
siz[i] = pp[i].n[ij[i]++];
                                                          while (ni[i] = pp[i].x[ij[i]])
                                                                   siz[i] + = pp[i].n[ij[i] + +];
}
                                                ni[i]++;
                                       }
                              if (++nn = = olen | ] !more && nn) {
   45
                                       dumpblock();
                                       for (i = 0; i < 2; i++)
                                                po[i] = out[i];
                                       nn = 0;
                              }
   50
            * dump a block of lines, including numbers, stars: pr_align()
   55
            */
            static
                                                                                                                 dumpblock
            dumpblock()
                     register i;
   60
                     for (i = 0; i < 2; i++)
                                                                                               Page 4 of nwprint.c
                              *po[i] - = '0';
```

### FIGURE 4K

```
...dumpblock
                     (void) putc('\n', fx);
     5
                     for (i = 0; i < 2; i++) {
                              if (*out[i] && (*out[i] != ' ' | | *(po[i]) != ' ')) {
                                       if (i = 0)
                                                 nums(i);
                                       if (i == 0 && *out[1])
   10
                                                 stars();
                                       putline(i);
                                       if (i == 0 \&\& *out[1])
                                                 fprintf(fx, star);
                                        if (i = = 1)
   15
                                                 nums(i);
                              }
                     }
            }
    20
* put out a number line: dumpblock()
            static
                                                                                                                          nums
            nums(ix)
    25
                                        /* index in out[] holding seq line */
                     int
            {
                                        nline[P_LINE];
                     char
                     register
TU
                                        *pn, *px, *py;
                     register char
    30
                     for (pn = nline, i = 0; i < lmax + P_SPC; i++, pn++)
                               *pn = ' ';
                     for (i = nc[ix], py = out[ix]; *py; py++, pn++) {
                              if (*py == ' ' || *py == '-')
35
                                        *pn = .
                               else {
                                        if (i\%10 = 0) || (i = 1 \&\& nc[ix]! = 1) {
                                                 j = (i < 0)? -i : i;
                                                 for (px = pn; j; j /= 10, px--)
    40
                                                           px = j\%10 + '0';
                                                 if (i < 0)
                                                           *px = '-';
                                        }
                                        else
    45
                                                 *pn = ' ';
                                        i++;
                               }
                     *pn = '\0';
    50
                     nc\{ix\} = i;
                     for (pn = nline; *pn; pn++)
                             (void) putc(*pn, fx);
                     (void) putc('\n', fx);
    55
             * put out a line (name, [num], seq, [num]): dumpblock()
            static
                                                                                                                        putline
            putline(ix)
    60
                               ix;
                                                                                      Page 5 of nwprint.c
```

### FIGURE 4L

```
...putline
                                        ī;
                     int
                                        *px;
      5
                     register char
                     for (px = namex[ix], i = 0; *px && *px != ':'; px++, i++)
                               (void) putc(*px, fx);
                     for (; i < lmax+P_SPC; i++)
                               (void) putc(' ', fx);
    10
                     /* these count from 1:
                      * ni∏ is current element (from 1)
                      * nc[] is number at start of current line
    15
                     for (px = out[ix]; *px; px++)
                               (void) putc(*px&0x7F, fx);
                      (void) putc('\n', fx);
            }
    20
* put a line of stars (seqs always in out[0], out[1]): dumpblock()
     25
             static
                                                                                                                            stars
             stars()
<u>}</u>=
Ĺ
                      register char
                                         *p0, *p1, ex, *px;
     30
                      if (!*out[0] || (*out[0] == ' ' && *(po[0]) == ' ') ||
                         !*out[1] | [ (*out[1] == ``&& *(po[1]) == ``))
                               return;
                      px = star;
     35
                      for (i = Imax + P_SPC; i; i--)
^{*}px++='';
                      for (p0 = out[0], p1 = out[1]; *p0 && *p1; p0++, p1++) {
ļ.
                                if (isalpha(*p0) && isalpha(*p1)) {
     40
                                         if (xbm[*p0-'A']&xbm[*p1-'A']) {
                                                  cx = \frac{1}{2};
                                                   nm + +;
                                         else if (!dna && _day[*p0-'A'][*p1-'A'] > 0)
     45
                                                   \epsilon x = '.\bar{};
                                         else
                                                   cx = ' ;
                                }
     50
                                else
                                         cx =
                                *px++=cx;
                       *px++ = '\n';
                       *px = '0';
     55
```

### FIGURE 4M

```
* strip path or prefix from pn, return len: pr_align()
            */
     5
           static
                                                                                                            stripname
           stripname(pn)
                                     /* file name (may be path) */
                    char
                             *pn;
                    register char
                                     *px, *py;
    10
                    py = 0;
                    for (px = pn; *px; px++)
                             if (*px = = T)
                                     py = px + 1;
    15
                    if (py)
                             (void) strcpy(pn, py);
                   . return(strlen(pn));
           }
    20
40
    45
    50
    55
```

60

### FIGURE 4N

```
* cleanup() - cleanup any ump file
             * getseq() -- read in seq, set dna, len, maxlen
     5
             * g calloc() -- calloc() with error checkin
             * readjmps() -- get the good jmps, from tmp file if necessary
             * writejmps() -- write a filled array of jmps to a tmp file: nw()
             */
            #include "nw.h"
   10
            #include < sys/file.h>
                      *jname = "/tmp/homgXXXXXX";
                                                                      /* tmp file for imps */
            FILE
                                                                      /* cleanup tmp file */
   15
                      cleanup();
            int
            long
                      lseek();
             * remove any tmp file if we blow
    20
                                                                                                                           cleanup
            cleanup(i)
if (fj)
   25
                                (void) unlink(jname);
7
                      exit(i);
ļå
ſΩ
ļá
             * read, return pir to seq, set dna, len, maxlen
    30
fŲ
             * skip lines starting with ';', '<', or '>'
             * seq in upper or lower case
E
             */
char
                                                                                                                             getseq
    35
             getseq(file, len)
                                          /* file name */
                                *file;
                      char
                                *len;
                                          /* seq len */
                      int
<u>ļ</u>å
                                          line[1024], *pseq;
                      char
    40
                      register char
                                          *px, *py;
                      int
                                          natge, tlen;
                      FILE
                                          *fp;
                      if ((fp = fopen(file, "r")) == 0) {
    45
                                fprintf(stderr," %s: can't read %s\n", prog, file);
                                exit(1);
                      tlen = natgc = 0;
                       while (fgets(line, 1024, fp)) {
                                if (*line == ';' | | *line == '<' | | *line == '>')
    50
                                for (px = line; *px != '\n'; px++)
                                          if (isupper(*px) || islower(*px))
                                                   tlen++;
    55
                       if ((pseq = malloc((unsigned)(llen+6))) = = 0) {
                                fprintf(stderr," %s: malloc() failed to get %d bytes for %s\n", prog, tlen+6, file);
                                exit(1);
    60
                       pseq[0] = pseq[1] = pseq[2] = pseq[3] = '\0';
```

```
...getseq
                     py = pseq + 4;
                     *len = tlen;
     5
                     rewind(fp);
                     while (fgets(line, 1024, fp)) {
                              if (*line == ';' || *line == '<' || *line == '>')
                                       continue;
   10
                              for (px = line; *px != '\n'; px + +) {
                                       if (isupper(*px))
                                                *py++ = *px;
                                       else if (islower(*px))
                                                 *py++ = toupper(*px);
                                       if (index("ATGCU",*(py-1)))
   15
                                                natge++;
                              }
                     *py++ = '\0';
   20
                     *py = '\0';
(void) fclose(fp);
                     dna = natgc > (tlen/3);
                     return(pseq+4);
            }
   25
10
            char
                                                                                                                     g_calloc
į.
            g_calloc(msg, nx, sz)
                                                 /* program, calling routine */
Щ
                                                /* number and size of elements */
                     int
    30
*px, *calloc();
                     if ((px = calloc((unsigned)nx, (unsigned)sz)) = = 0) {
                              if (*msg) {
                                        fprintf(stderr, "%s: g_calloc() failed %s (n=%d, sz=%d)\n", prog, msg, nx, sz);
    35
                                       exit(1);
                              }
                     return(px);
    40
            }
             * get final jmps from dx[] or tmp file, set pp[], reset dmax: main()
                                                                                                                   readjmps
    45
            readjmps()
                                       fd = -1;
                                        siz, i0, i1;
                     register i, j, xx;
    50
                     if (fj) {
                              (void) fclose(fj);
                              if ((fd = open(jname, O_RDONLY, 0)) < 0) {
                                        fprintf(stderr, "%s: can't open() %s\n", prog, jname);
    55
                                        cleanup(1);
                              }
                     for (i = i0 = i1 = 0, dmax0 = dmax, xx = len0; ; i++) {
                              while (1) {
                                        for (j = dx[dmax].ijmp; j > = 0 && dx[dmax].jp.x[j] > = xx; j-)
    60
                                                                                               Page 2 of nwsubr.c
```

### FIGURE 4P

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<u>‡</u> 30

35

40

45

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60

} else

if (fd > = 0)

if (fj) {

(void) close(fd);

offset = 0;}}

 $f_1 = 0$ ;

(void) unlink(jname);

...readjmps if  $(j < 0 && dx[dmax].offset && fj) {$ (void) Iseek(fd, dx[dmax].offset, 0); (void) read(fd, (char \*)&dx[dmax].jp, sizeof(struct jmp)); (void) read(fd, (char \*)&dx[dmax].offset, sizeof(dx[dmax].offset)); dx[dmax].ijmp = MAXJMP-1;} else break; if (i > = JMPS) { fprintf(stderr, "%s: too many gaps in alignment\n", prog); cleanup(1); if (j. > -0) { siz = dx[dmax].jp.n[j];xx = dx[dmax].jp.x[j];dmax += siz;if (siz < 0) { /\* gap in second seq \*/ pp[1].n[i1] = -siz;xx + = siz;/\* id = xx - yy + len1 - 1pp[1].x[i1] = xx - dmax + len1 - 1;gapy + +;ngapy -= siz; /\* ignore MAXGAP when doing endgaps \*/ siz = (-siz < MAXGAP | | endgaps)? -siz : MAXGAP; 11++; else if (siz > 0) { /\* gap in first seq \*/ pp[0].n[i0] = siz;pp[0].x[i0] = xx;gapx + +;ngapx += siz; /\* ignore MAXGAP when doing endgaps \*/ siz = (siz < MAXGAP | endgaps)? siz : MAXGAP; } break; /\* reverse the order of jmps for (j = 0, i0--; j < i0, j++, i0--)i = pp[0].n[j]; pp[0].n[j] = pp[0].n[i0]; pp[0].n[i0] = i;i = pp[0].x[j]; pp[0].x[j] = pp[0].x[i0]; pp[0].x[i0] = i;for (j = 0, i1--; j < i1; j++, i1--)i = pp[1].n[j]; pp[1].n[j] = pp[1].n[i1]; pp[1].n[i1] = i;i = pp[1].x[j]; pp[1].x[j] = pp[1].x[i1]; pp[1].x[i1] = i;

### FIGURE 4Q

```
* write a filled jmp struct offset of the prev one (if any): nw()
     5
            */
                                                                                                                     writejmps
            writejmps(ix)
                     int
                               ix;
            {
                               *mktemp();
                     char
   10
                     if (!fj) {
                               if (mktemp(jname) < 0) {
                                        fprintf(stderr, "%s: can't mktemp() %s\n", prog, jname);
    15
                               if ((f_j - fopen(jname, "w")) = = 0) {
                                         fprintf(stderr, "%s: can't write %s\n", prog, jname);
                               }
    20
                     (void) fwrite((char *)&dx[ix].jp, sizeof(struct jmp), 1, fj);
                     (void) fwrite((char *)&dx[ix].offset, sizeof(dx[ix].offset), 1, fj);
    25
30
10
L4
fIJ
    35
ļi
1 40
1 40
    45
    50
    55
    60
```

GTGCTCTCCGAGGACAAGCAGGAGGNGGTGGAGCTGGTGAAGCACCATCTGTGGGCTCTG
GAAGTGTGCTACATCTCAGCCTTGGTCTTGTCCTGCTTACTCACCTTCCTGGTCCTGATG
CGCTCACTGGTGACACACAGGACCAACCTTCGAGCTCTGCACCGAGGAGCTGCCCTGGAC
TTGAGTCCCTTGCATCGGAGTCCCCATCCCTCCCGCCAAGCCATATTCTGTTGGATGAGC

1 0 TTCAGTGCCTACCAGACAGCCTTTATCTGCCTTGGGCTCCTGGTGCAGCAGATCATCTTC
TTCCTGGGAACCACGGCCTTGGCCTTCCTGGTGCTCCTGTGCTCCATGGCAGGAAC
CTCCTGCTCTTCCGTTCCCTGGAGTCCTCGTGGCCCTTCTGGCTGACTTTGGCCCTGGCT
GTGATCCTGCAGAACATGGCAGCCCATTGGGTCTCCTGGAGACTCATGATGGACACCCA
CAGCTGACCAACCGGCGAGTGCTCTATGCAGCCACCTTTCTTCTCTCCCCCTCAATGTG
15 CTGGTGGGTGCCATGGTGGCCACCTGGCGAGAGCCGCCACTCTCGACCCCGC
TACTACACGTACCGAA

CACAACCAGCCACCCTCTAGGATCCCAGCCCAGCTGGTGCTGGGCTCAGAGGAAGGC 5 CCCGTGTTGGGAGCACCCTGCTTGCCTGGAGGGACAAGTTTCCGGGAGAGATCAATAAAG GAAAGGAAAGACAAGGAAGGGAGAGGTCAGGAGAGCGCTTGATTGGAGGAGAAGGGCC AGAGAATGTCGTCCCAGCCAGCAGGGAACCAGACCTCCCCCGGGGCCACAGAGGACTACT 10 TGTCAATCCTTGTGCTGCTGCTCCTGGCCATGCTGAGGCGCCGCCAGCTCTGGCCTG ACTGTGTGCGTGGCAGGCCCGGCCTGCCCAGGCCCCGGGCAGTGCCTGCTGTTTTCA TGGTCCTCCTGAGCTCCCTGTGTTTGCTGCTCCCCGACGAGGACGCATTGCCCTTCCTGA CTCTCGCCTCAGCACCCAGCCAAGATGGGAAAACTGAGGCTCCAAGAGGGGCCTGGAAGA TACTGGGACTGTTCTATTATGCTGCCCTCTACTACCCTCTGGCTGCCTGTGCCACGGCTG 15 GCCACACAGCTGCACACCTGCTCGGCAGCACGCTGTCCTGGGCCCACCTTGGGGTCCAGG TCTGGCAGAGGGCAGAGTGTCCCCAGGTGCCCAAGATCTACAAGTACTACTCCCTGCTGG CCTCCCTGCCTCCCTGCTGGGCCTCGGATTCCTGAGCCTTTGGTACCCTGTGCAGCTGG TGAGAAGCTTCAGCCGTAGGACAGGAGCAGGCTCCAAGGGGCTGCAGAGCAGCTACTCTG AGGAATATCTGAGGAACCTCCTTTGCAGGAAGAAGCTGGGAAGCAGCTACCACACCTCCA 20 AGCATGGCTTCCTGTCCTGGGCCCGCGTCTGCTTGAGACACTGCATCTACACTCCACAGC CAGGATTCCATCTCCCGCTGAAGCTGGTGCTTTCAGCTACACTGACAGGGACGGCCATTT ACCAGGTGGCCCTGCTGCTGGTGGGCGTGGTACCCACTATCCAGAAGGTGAGGGCAG GGGTCACCACGGATGTCTCCTACCTGCTGGCCGGCTTTGGAATCGTGCTCTCCGAGGACA AGCAGGAGGTGGTGGAGCTGGTGAAGCACCATCTGTGGGCTCTGGAAGTGTGCTACATCT 25 CAGCCTTGGTCTTGTCCTGCTTACTCACCTTCCTGGTCCTGATGCGCTCACTGGTGACAC ACAGGACCAACCTTCGAGCTCTGCACCGAGGAGCTGCCCTTGGACTTGAGTCCCTTGCATC GGAGTCCCCATCCCTCCCGCCAAGCCATATTCTGTTGGATGAGCTTCAGTGCCTACCAGA CAGCCTTTATCTGCCTTGGGCTCCTGGTGCAGCAGATCATCTTCTTCCTGGGAACCACGG CCCTGGCCTTCCTGGTGCTCATGCCTGTGCTCCATGGCAGGAACCTCCTGCTCTTCCGTT 30 CCCTGGAGTCCTCGTGGCCCTTCTGGCTGACTTTGGCCCTGGCTGTGATCCTGCAGAACA TGGCAGCCCATTGGGTCTTCCTGGAGACTCATGATGGACACCCACAGCTGACCAACCGGC TGGCCACCTGGCGAGTGCTCCTCTCTGCCCTCTACAACGCCATCCACCTTGGCCAGATGG ACCTCAGCCTGCCGCCACCGAGAGCCGCCACTCTCGACCCCGGCTACTACACGTACCGAA 35 ACTTCTTGAAGATTGAAGTCAGCCAGTCGCATCCAGCCATGACAGCCTTCTGCTCCCTGC TCCTGCAAGCGCAGAGCCTCCTACCCAGGACCATGGCAGCCCCCCAGGACAGCCTCAGAC CAGGGGAGGAAGACGAAGGGATGCAGCTGCTACAGACAAAGGACTCCATGGCCAAGGGAG CTAGGCCCGGGGCCAGCCGCGGCAGGGCTCGCTGGGGTCTGGCCTACACGCTGCTGCACA ACCCAACCCTGCAGGTCTTCCGCAAGACGCCCTGTTGGGTGCCAATGGTGCCCAGCCC**T** 40 **GA**GGGCAGGGAAGGTCAACCCACCTGCCCATCTGTGCTGAGGCATGTTCCTGCCTACCAC GGATCACTGTGGTTGGGTGGAGGTCTGTCTGCACTGGGAGCCTCAGGAGGGCTCTGCTCC CCTTGGTCCAGGAGCCAGTTGAGCCAGGGCAGCCACATCCAGGCGTCTCCCTACCCTGGC 45 TCTGCCATCAGCCTTGAAGGGCCTCGATGAAGCCTTCTCTGGAACCACTCCAGCCCAGCT CCACCTCAGCCTTGGCCTTCACGCTGTGGAAGCAGCCAAGGCACTTCCTCACCCCCTCAG CGCCACGGACCTCTCTGGGGAGTGGCCGGAAAGCTCCCGGGCCTCTGGCCTGCAGGGCAG CCCAAGTCATGACTCAGACCAGGTCCCACACTGAGCTGCCCACACTCGAGAGCCAGATAT TTTTGTAGTTTTTATGCCTTTGGCTATTATGAAAGAGGTTAGTGTGTTCCCTGCAATAAA 50 CTTGTTCCTGAGAAAAA

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MSSQPAGNQTSPGATEDYSYGSWYIDEPQGGEELQPEGEVPSCHTSIPPGLYHACLASL SILVLLLAMLVRRRQLWPDCVRGRPGLPRPRAVPAAVFMVLLSSLCLLLPDEDALPFL TLASAPSQDGKTEAPRGAWKILGLFYYAALYYPLAACATAGHTAAHLLGSTLSWAHLGV QVWQRAECPQVPKIYKYYSLLASLPLLLGLGFLSLWYPVQLVRSFSRRTGAGSKGLQSS YSEEYLRNLLCRKKLGSSYHTSKHGFLSWARVCLRHCIYTPQPGFHLPLKLVLSATLTG TAIYQVALLLLVGVVPTIQKVRAGVTTDVSYLLAGFGIVLSEDKQEVVELVKHHLWALE VCYISALVLSCLLTFLVLMRSLVTHRTNLRALHRGAALDLSPLHRSPHPSRQAIFCWMS FSAYQTAFICLGLLVQQIIFFLGTTALAFLVLMPVLHGRNLLLFRSLESSWPFWLTLAL AVILQNMAAHWVFLETHDGHPQLTNRRVLYAATFLLFPLNVLVGAIVATWRVLLSALYN AIHLGQMDLSLLPPRAATLDPGYYTYRNFLKIEVSQSHPAMTAFCSLLLQAQSLLPRTM AAPQDSLRPGEEDEGMQLLQTKDSMAKGARPGASRGRARWGLAYTLLHNPTLQVFRKTA LLGANGAQP

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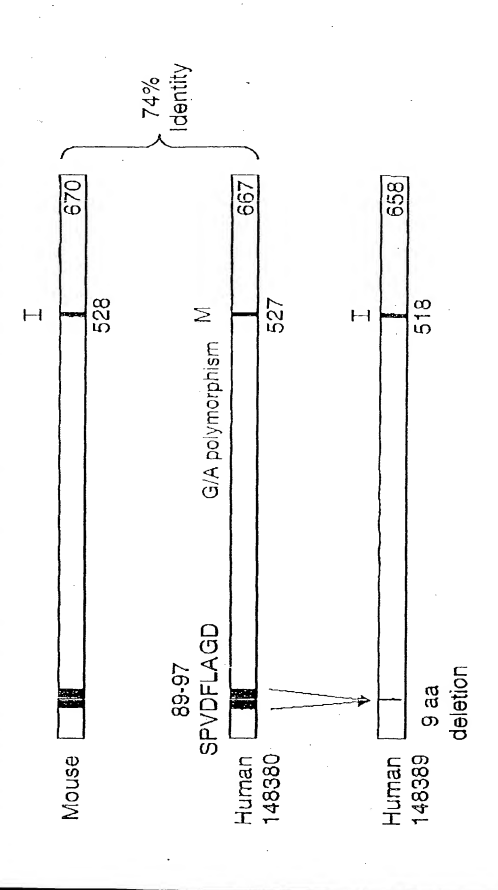
123-131

Important features of the protein:

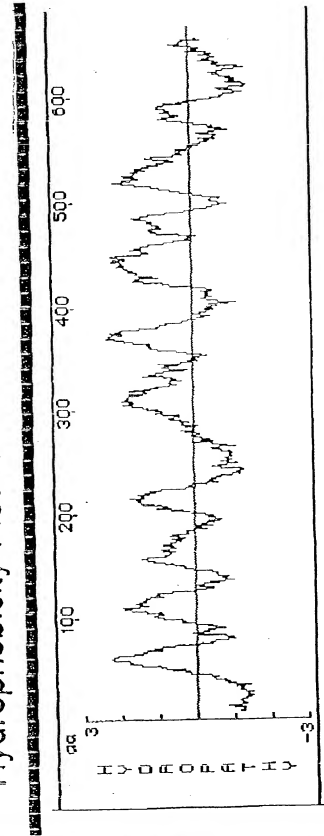
```
IJ
        Signal peptide:
none
  20
        Transmembrane domain:
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        54 - 71
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        93-111
14
        140-157
11 25
        197-214
        291-312
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[]
        356-371
        425-444
إية
        464 - 481
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  30
        505-522
Motif name: N-glycosylation site.
į.
               8 - 12
  35
        Motif name: N-myristoylation site.
             50-56
            167 - 173
  40
            232-238
            308 - 314
            332-338
            516-522
            618-624
  45
            622-628
            631-637
            652-658
        Motif name: Prokaryotic membrane lipoprotein lipid attachment
  50
        site.
            355-366
        Motif name: ATP/GTP-binding site motif A (P-loop).
```

## Stra6 Variant Clones

,是一个时间,我们的时间,我们就是一个时间,我们就是一个时间,我们是一个时间,我们也不是一个时间,我们是一个时间,我们也不是一个时间,我们也会会看到一个时间,我



## Hydrophobicity Plot of Human Stra6

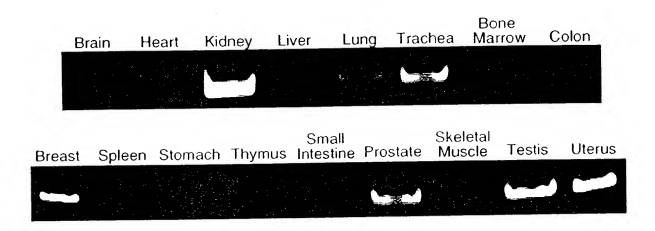


· 3 kb mRNA

667 Amino Acids -->50% Residues Hydrophobic

73.5 kDa Protein

9 Potential Transmembrane Domains



# Stra6 RNA Expression in Human Colon Tumor Tissue

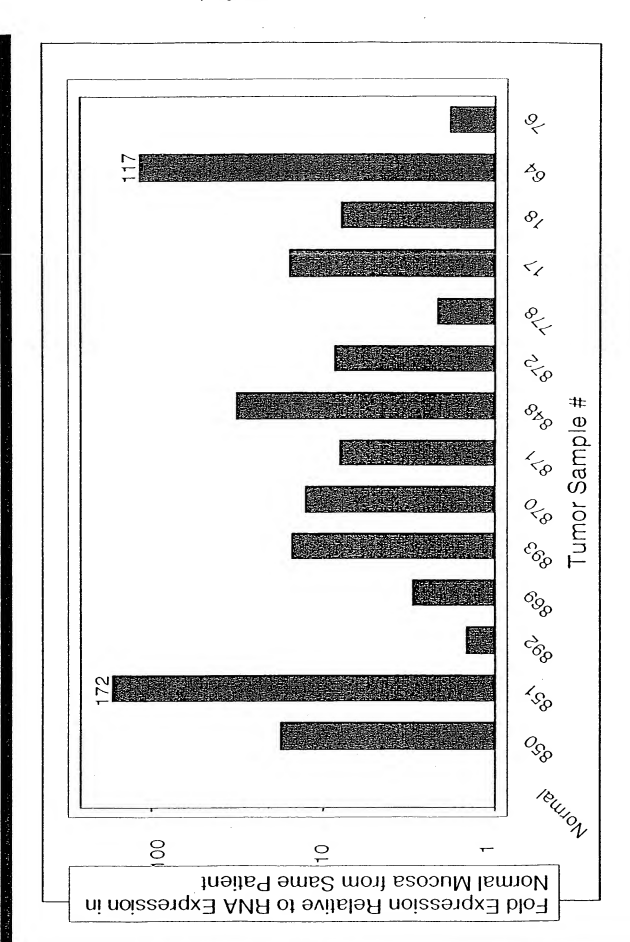
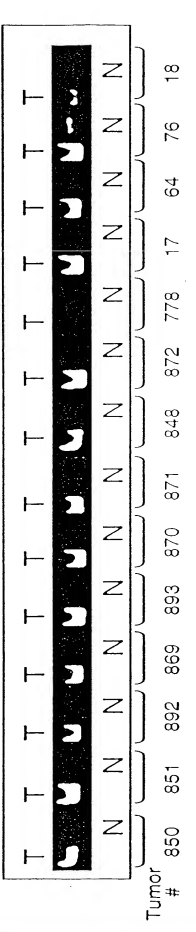


FIGURE 11

### issue vs Normal Mucosa From the Same Patient Stra6 RNA Expression in Human Colon Tumor

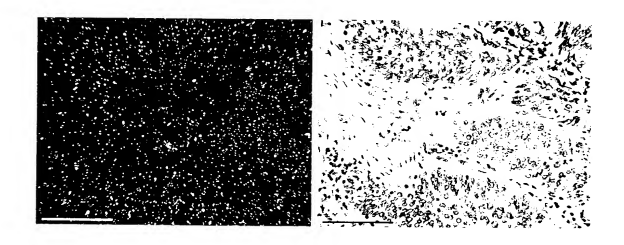
Taqman Product Analysis After 40 Cycles

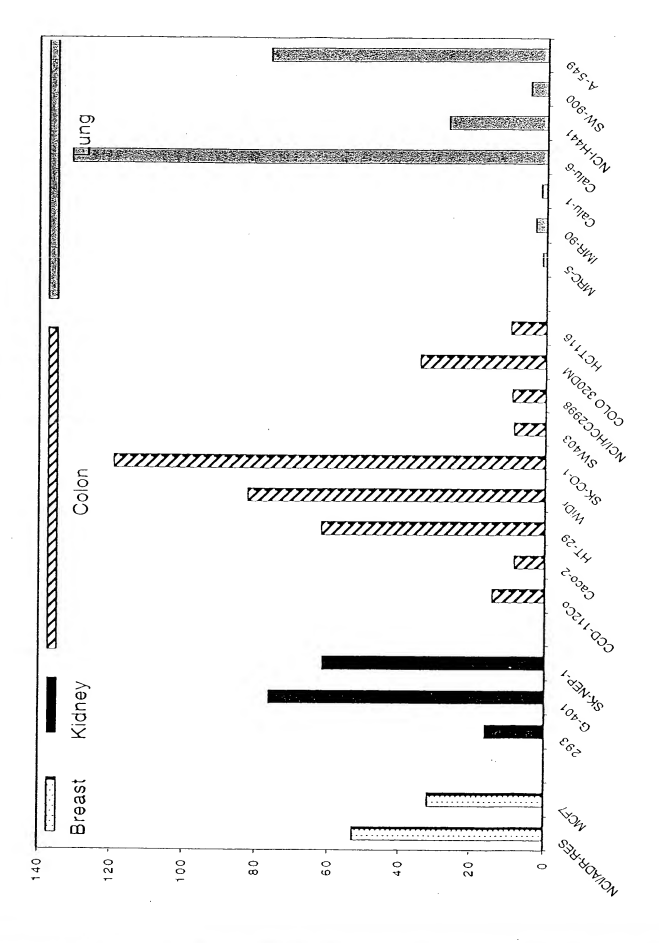
### Stra6



GAPDH

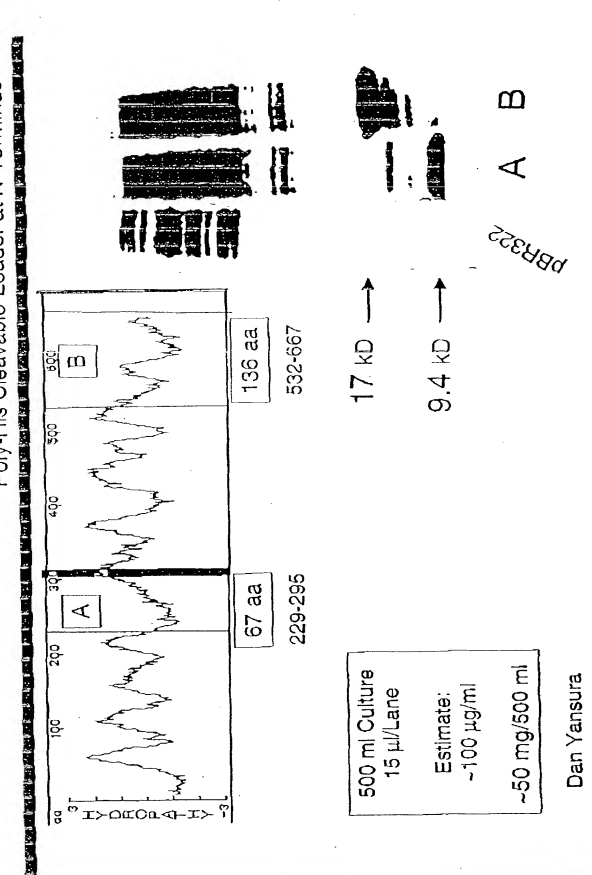






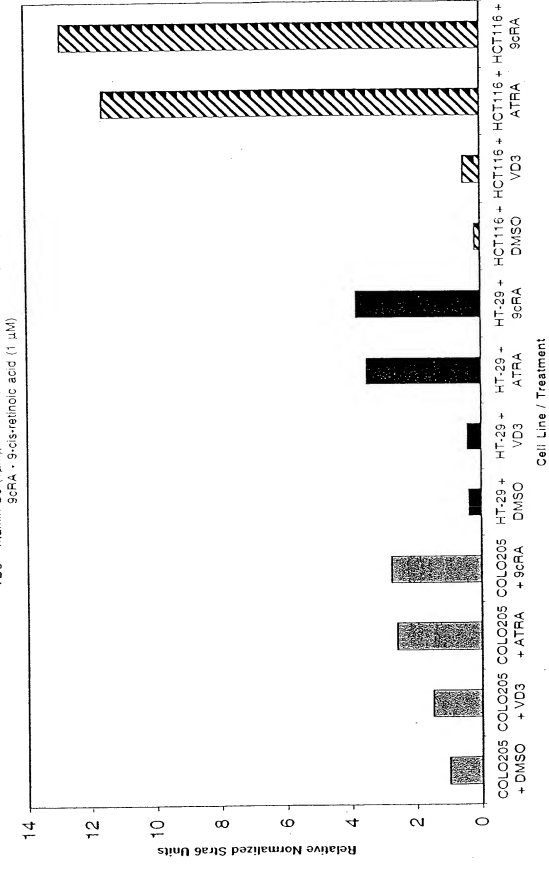
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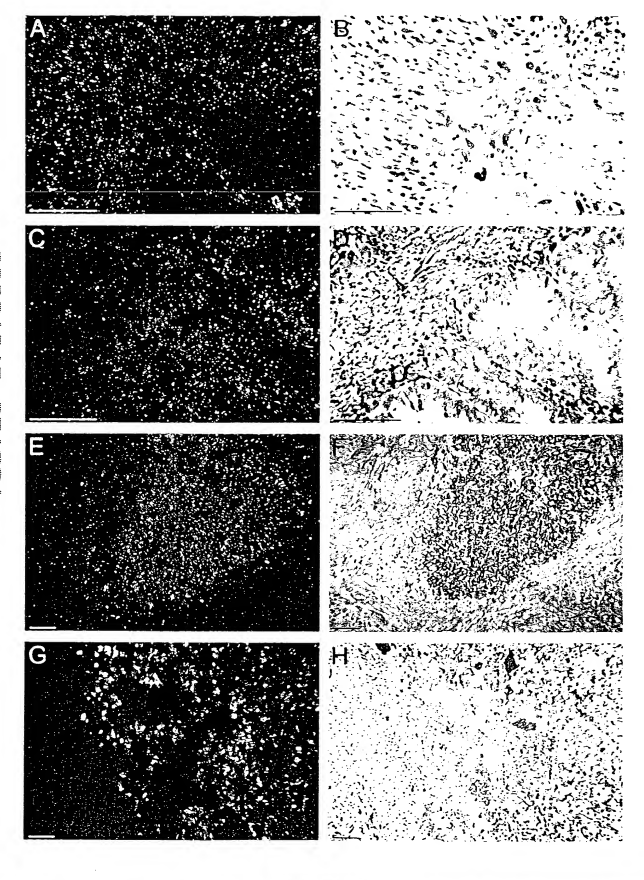
Poly-His Cleavable Leader at N-Terminus Stra6 Peptide Expression in E. coli



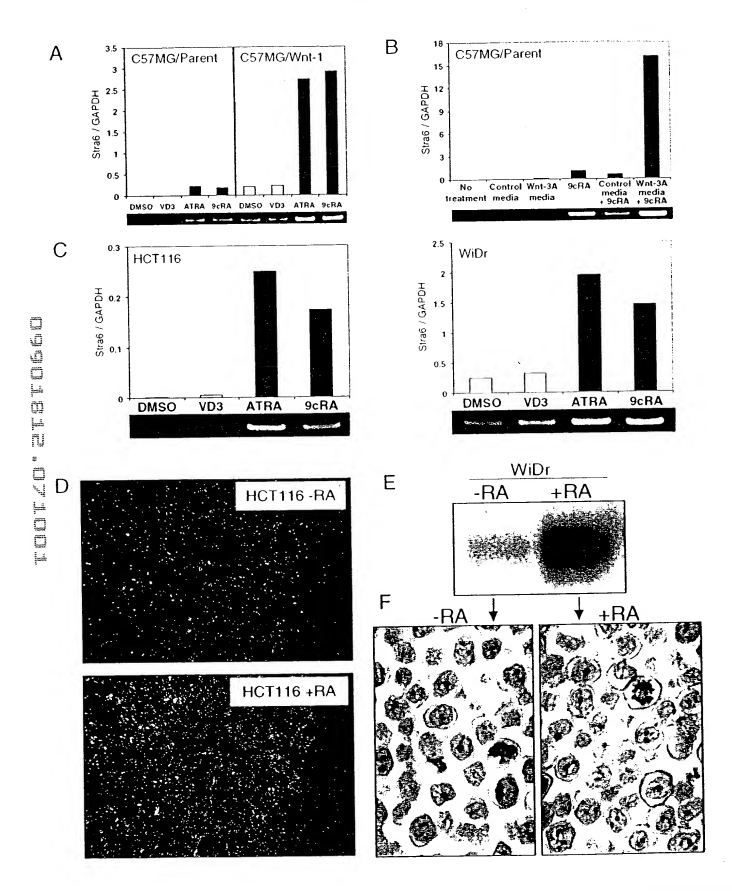
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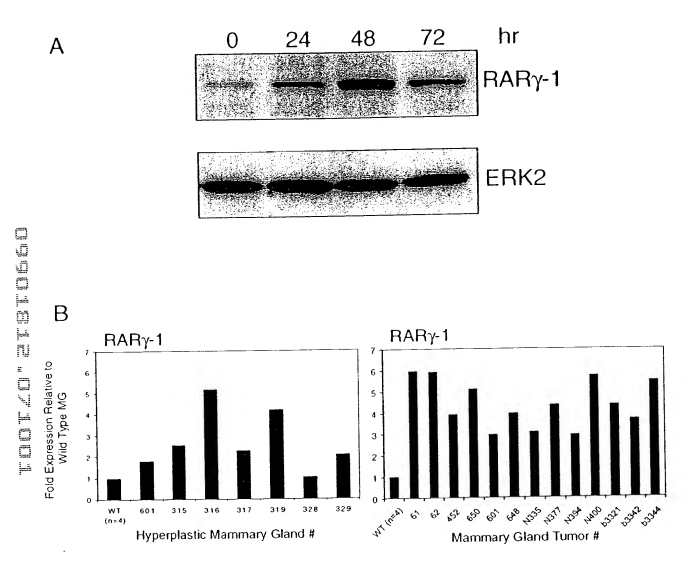
## Stra6 RNA Expression in Human Colon Carcinoma Cells +/- Retinoic Acid TM#75 (2/28/00) VD3 · vitamin D3 (1μΜ); ATRA · all·trans-retinoic acid (1 μΜ) 9cRA · 9-cis-retinoic acid (1 μΜ)





nggniale.cytoni





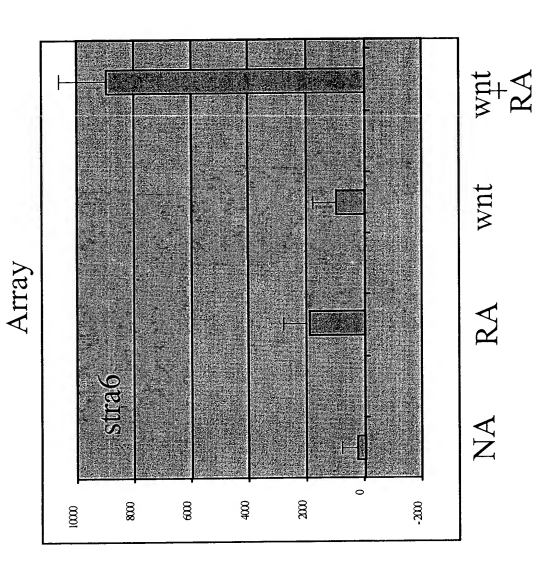
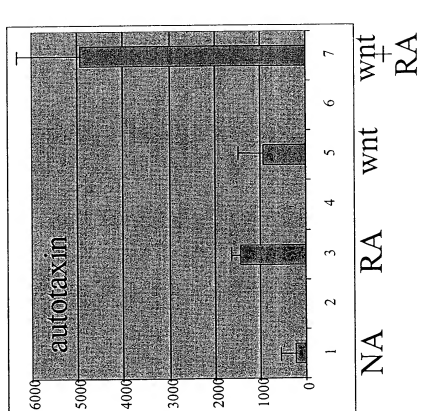


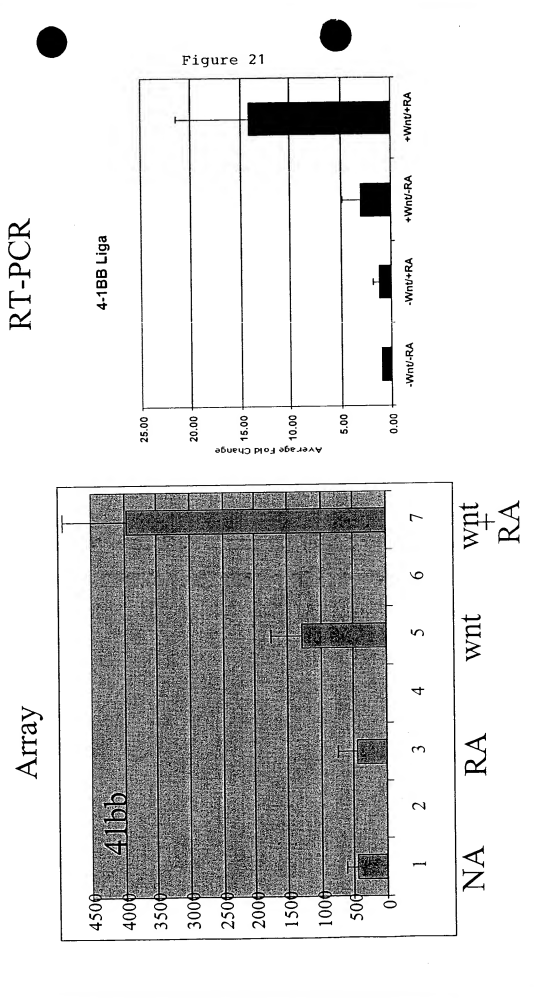
Figure 20

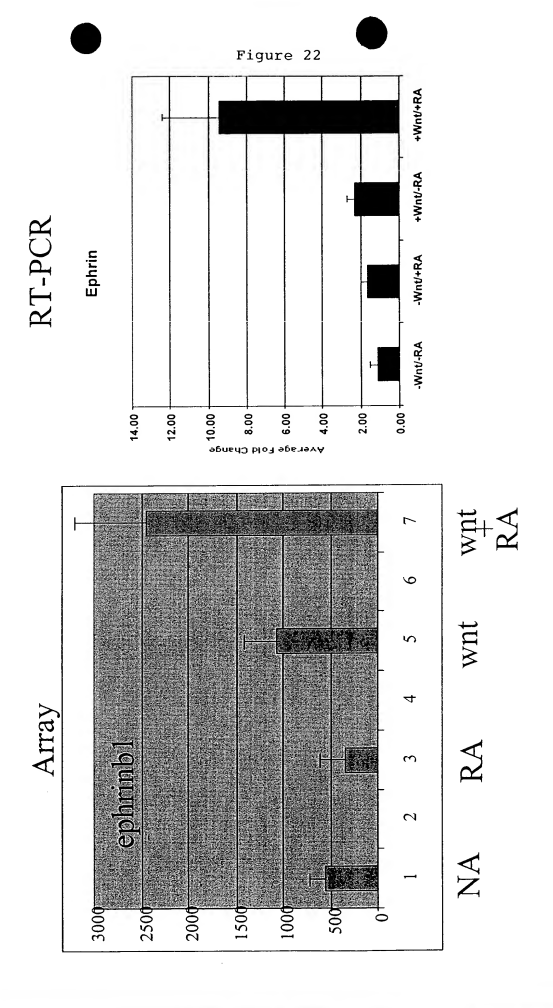
RT-PCR

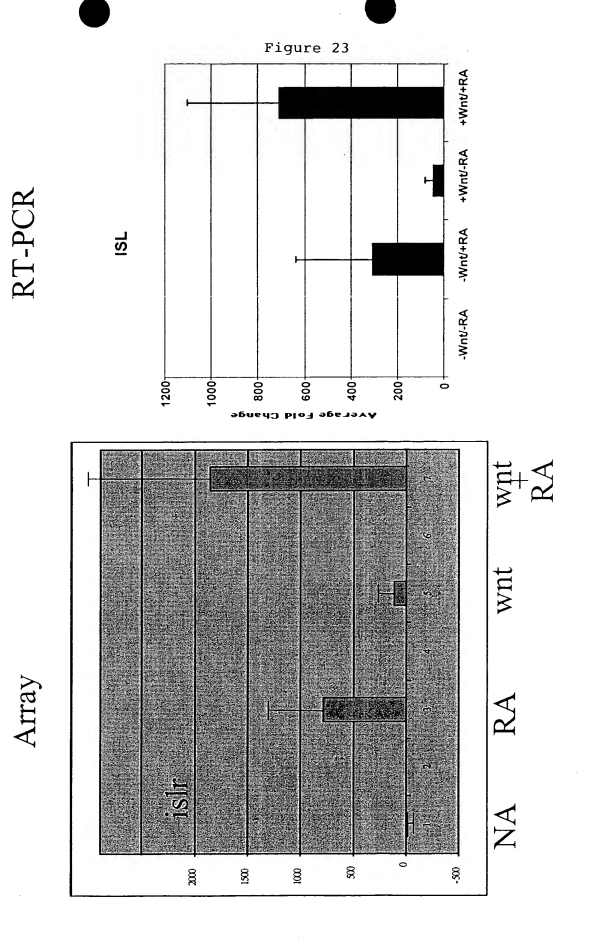
Array

+Wnt/+RA +Wnt/-RA Autotaxin -Wnt/+RA 30 2 9 20 6 9 Average Fold Change









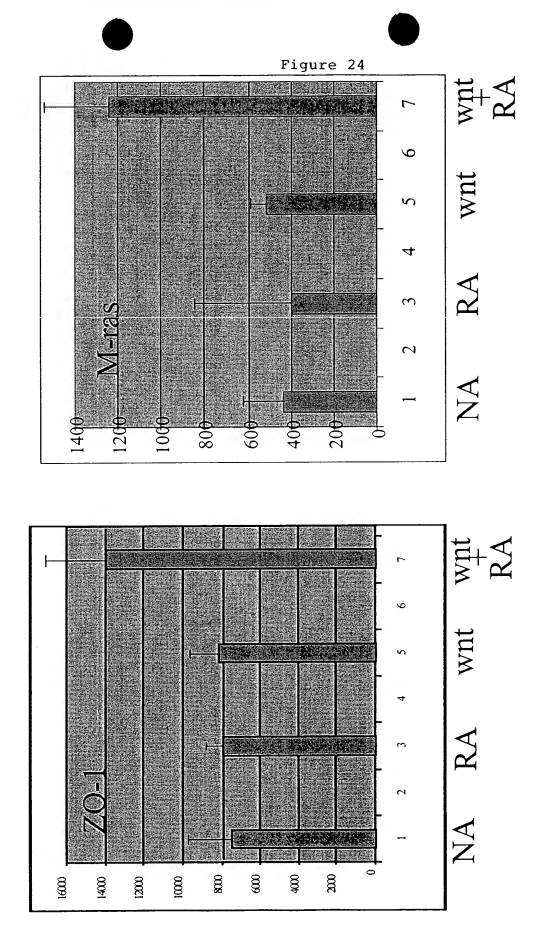
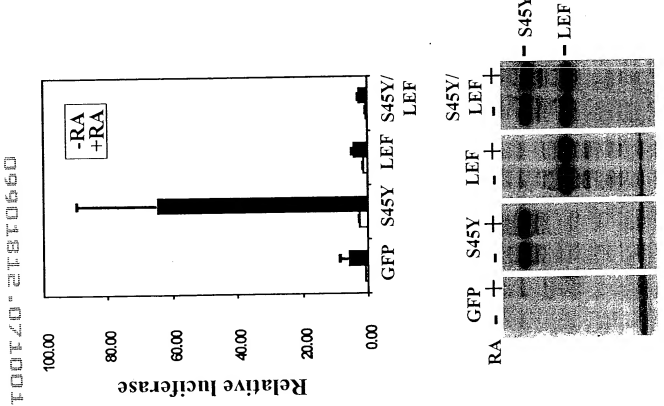


Figure 25

Figure 25 A

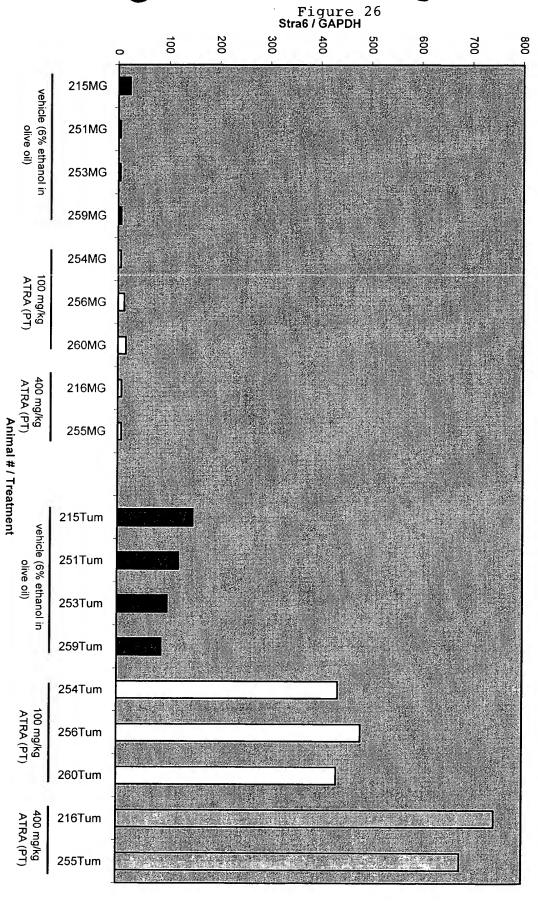
Figure 25 B



## Stra6 mRNA in normal mouse mammary gland and Wnt-1mammary gland tumors

Nude mice bearing Wnt-1 tumor explants were given ATRA peri-tumorally (PT) at 100 and 400 mg/kg.

Tumors and normal adjacent mammary glands were harvested 8 hours later.

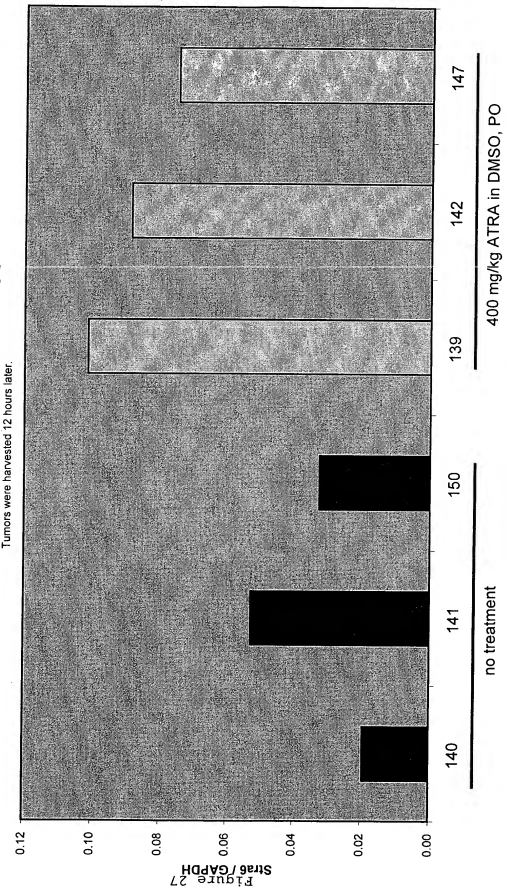


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### SCANNED, # 24

## Stra6 mRNA in WiDr xenografts from mice dosed with ATRA (400 mg/kg)

Nude mice bearing WiDr xenografts were given ATRA per orum (PO) at 400 mg/kg. Tumors were harvested 12 hours later.



Animal # / Treatment